Anaerobic methane oxidation driven by microbial reduction of natural organic matter in a tropical wetland

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Abstract

Wetlands constitute the main natural source of methane on Earth due to their high content of natural organic matter (NOM), but key drivers such as electron acceptors supporting methanotrophic activities in these habitats are poorly understood. We performed anoxic incubations using freshly collected sediment along with water samples harvested from a tropical wetland, amended with $^{13}$C-methane (0.67 atm) to test the capacity of its microbial community to perform anaerobic methane oxidation (AOM) linked to the reduction of the humic fraction of its NOM. Collected evidence demonstrates that electron-accepting functional groups (e.g. quinones) present in NOM fueled AOM by serving as terminal electron acceptor. Indeed, while sulfate reduction was the predominant process accounting for up to 42.5% of the AOM activities, microbial reduction of NOM concomitantly occurred. Furthermore, enrichment of wetland sediment with external NOM provided complementary electron-accepting capacity, which reduction accounted for $\sim 100$ nmol $^{13}$C-$\text{CH}_4$ oxidized cm$^{-3}$ d$^{-1}$. Spectroscopic evidence showed that quinone moieties were heterogeneously distributed in the wetland sediment, and that their reduction occurred during the course of AOM. Moreover, an enrichment derived from wetland sediments performing AOM linked to NOM reduction stoichiometrically oxidized methane coupled to the reduction of the humic analogue, anthraquinone-2,6-disulfonate. Microbial populations potentially involved in AOM coupled to microbial reduction of NOM were dominated by
divergent biota from putative AOM-associated archaea. We estimate that this microbial process could potentially contribute to the suppression of up to 114 Tg CH$_4$ yr$^{-1}$ in coastal wetlands and more than 1,300 Tg yr$^{-1}$ considering the global wetland area.

**Importance**

Identifying key processes governing methane emissions from natural systems is of major importance considering the global warming effects triggered by this greenhouse gas. AOM coupled to the microbial reduction of distinct electron acceptors plays a pivotal role in mitigating methane emissions from ecosystems. Given their high organic content, wetlands constitute the largest natural source of atmospheric methane. Nevertheless, processes controlling methane emissions in these environments are poorly understood. Here we provide tracer analysis with $^{13}$CH$_4$ and spectroscopic evidence revealing that AOM linked to the microbial reduction of redox functional groups in natural organic matter (NOM) prevails in a tropical wetland. We suggest that microbial reduction of NOM may largely contribute to suppress methane emissions from tropical wetlands. This is a novel avenue within the carbon cycle in which slowly decaying NOM (e.g. humic fraction) in organotrophic environments fuels AOM by serving as terminal electron acceptor.
Introduction

Microbial processes produce and consume methane (CH4) in anoxic sediments playing a crucial role in regulating Earth’s climate. Virtually 90% of CH4 produced from marine environments is oxidized by microorganisms avoiding its release to the atmosphere (1).

Anaerobic oxidation of methane (AOM) associated with sulfate reduction was first discovered in marine environments (2). More recently, AOM has also been linked to the microbial reduction of nitrate (3, 4) and nitrite (5), as well as Fe(III) and Mn(IV) oxides (6–8) in freshwater and marine environments. Wetlands are the largest natural source of CH4 (9), contributing to about a third of global emissions (10), but key drivers, such as electron acceptors fueling methanotrophic activities in these habitats, are poorly understood. CH4 emissions from wetlands have been strongly responsive to climate in the past, and will likely continue to be responsive to anthropogenic-driven climate change in the future, predicting a large impact on global atmospheric CH4 concentration (10). The traditional assumption is that aerobic methanotrophy dominates wetlands’ CH4 cycling by oxidizing an estimated 40 to 70% of gross CH4 production in these ecosystems (11). Recent findings (12) challenged this conjecture by providing evidence that AOM may consume up to 200 Tg CH4 yr⁻¹, decreasing their potential CH4 emission by 50% in these habitats. Most AOM activities observed in wetlands have been related to sulfate reduction (12, 13), but other electron acceptors remain feasible. Natural organic matter (NOM), circumscribed to humic substances (HS) in many studies (14), occurs at high concentrations in wetlands both in soluble and solid phases (15). Recent evidence indicates that HS suppress methane production in different ecosystems (16, 17), yet the mechanisms involved are still enigmatic. HS can theoretically promote AOM as they can serve as terminal electron acceptors.
acceptors for microbial respiration (18, 19) and have higher redox potential than sulfate (20). However, compelling evidence demonstrating AOM driven by the microbial reduction of NOM present in anoxic environments remains elusive (21, 22).

We aimed to document $^{13}$CH$_4$ anaerobic oxidation and the ongoing reduction of intrinsic electron acceptors, including the electron accepting fraction of NOM, by the biota of freshly sampled sediment from a coastal tropical wetland. We provide $^{13}$CH$_4$ tracer studies and spectroscopic evidence demonstrating for the first time that AOM is linked to the microbial reduction of redox functional groups present in the NOM of this tropical marsh. Furthermore, we found evidence, based on 16S rRNA gene sequences, indicating that microbial populations potentially involved in AOM coupled to microbial reduction of NOM were dominated by divergent biota from putative AOM-associated microorganisms.

**Results**

**Kinetics of $^{13}$C-methane oxidation and electron balances**

Exponential phase of AOM was observed in microcosms over the first 15 days of incubation in the case of unamended sediment (free from external NOM addition). The methanotrophic rate in this experimental treatment was $\sim$1.34 $\mu$mol $^{13}$C-methane oxidized cm$^{-3}$ d$^{-1}$ (Fig. 1). At the end of the exponential phase, sulfate and Fe(III) reduction accounted for 42.5% and 0.5% of $^{13}$C-methane oxidized, respectively, while the role of nitrate was marginal (Fig. 2 and Table S2). These unamended sediment microcosms exhibited a reduction in intrinsic NOM during the course of AOM, which was expected due to the high concentration of organic carbon in the tropical wetland, with the capacity to
accept electrons (Table S1, Fig. 2). Nevertheless, large perturbation caused by endogenous NOM reduction in experimental controls lacking $^{13}$C-methane obstructed accurate assessment of AOM driven by this microbial process (Fig. 2). The large endogenous NOM reduction observed in these control experiments may be explained by concomitant methane production (and subsequent consumption) observed (Fig. S2), and by oxidation of labile organic matter present in the sediment (Table S1). Supplementary incubations spiked with the sulfate-reduction inhibitor, sodium molybdate (25 mM), showed decreased sulfate reducing activities (~50%, Fig. 2), while AOM rates remained high when compared against their non-inhibited counterparts (Fig. 1). Remarkably, when sulfate reduction was inhibited, the reduction of intrinsic NOM was doubled (from 1.6±0.11 to 3.4±0.19 milli-electron equivalents (meq) l$^{-1}$), implying that the reduction of redox functional groups in NOM was promoted when the utilization of sulfate was impeded.

Further enrichment of wetland sediment with external NOM, in the form of HS derived from Pahokee Peat (Florida Everglades, 2.5 g l$^{-1}$), provided complementary electron accepting capacity, which significantly elicited AOM up to ~1.88 µmol $^{13}$C-methane oxidized cm$^{-3}$ d$^{-1}$ and extended the exponential phase to 20 days (Fig. 1). In this experimental treatment, electron balances revealed a methanotrophic activity responsible of ~100 nmol $^{13}$C-CH$_4$ oxidized cm$^{-3}$ d$^{-1}$ linked to microbial reduction of NOM (including both intrinsic and externally added as Pahokee Peat HS). As hypothesized before, consumption of intrinsically produced methane was confirmed by experimental controls enriched with HS from Pahokee Peat and incubated in the absence of $^{13}$C-methane, which showed significant consumption of $^{12}$CH$_4$ (Fig. S2). This was also confirmed by increased $^{12}$CO$_2$ production quantified, which was reflected on 2 to 4-fold lower enrichment of $^{13}$CO$_2$. 
in HS enriched incubations as compared to unamended controls (see $^{13}$FCO$_2$ values in Fig. 1). Reports (23, 24) indicate that methanotrophic microorganisms prefer to oxidize $^{12}$CH$_4$ as compared to $^{13}$CH$_4$, which may partly explain our findings.

The role of sulfate reduction on AOM when wetland sediment was enriched with HS was not possible to assess (Table S2) due to large endogenous sulfate reduction elicited by degradation of the labile fraction of externally added NOM (Fig. 2), which also triggered methanogenesis in these microcosms. Since no significant differences in iron reduction were detected between microcosms with or without $^{13}$CH$_4$ addition, the only microbial process clearly identified driving AOM in Pahokee Peat enriched sediments was the microbial reduction of HS (Table S2).

**Spectroscopic evidence on presence and reduction of redox-functional groups in NOM**

Initial exploration of the solid phase NOM present in wetland sediment by micro-ATR-FTIR spectra, revealed the presence of electron accepting moieties both in unamended and in HS enriched wetland sediments. By mapping of acquisition points at 1650-1620 cm$^{-1}$, presence and heterogeneous distribution of quinone functional groups was evidenced in sediments confirming the presence of non-soluble electron accepting moieties classically attributed to humic-like materials (Fig. 3a and b). To further confirm this, we looked for double bonded carbon and oxygen (C=O) by use of X-ray photoelectron spectra (XPS), technique that supported the existence of quinone-like functional groups in unamended sediment and furthermore, provided evidence of the reduction of these moieties by showing the disappearance of the C=O signal from C1s and O1s high resolution spectra when comparing signals from sediment analyzed before and after incubation with $^{13}$CH$_4$ in the
absence of external HS (Fig. 3c to f). Another missing signal after the AOM process was that which corresponds to metallic oxides, evidenced by analysis of the O1s high resolution spectra (Figure 3d and f), which may imply reduction of intrinsic iron oxides that supported ~0.5% of methanotrophy according to electron balances (Table S2). Further analysis of the liquid phase of pristine sediment microcosms also revealed the reduction of quinone-like moieties during the course of AOM (Fig. 4). Initial samples exhibited a well-defined and strong peak at 1690 cm\(^{-1}\) associated with quinone moieties, while reduced samples, at the end of the incubation period, showed an increase in the signal related to phenolic groups (1660 cm\(^{-1}\)). Additional signals of phenolic groups were detected after incubation with \(^{13}\)CH\(_4\) and Pahokee Peat by spectral signals detected around 2260-2500 cm\(^{-1}\) (25).

**Microbial communities performing AOM**

According to 16S rRNA gene sequences from wetland sediment samples performing AOM, anaerobic methanotrophic archaea (ANME), which are traditionally linked to anaerobic methanotrophy under sulfate-reducing (2, 26), Fe(III)-reducing (6, 8), and artificial electron acceptor-reducing conditions (27), were barely detected in our experiments, with ANME-1b and ANME-3 representing less than 0.5% and 0.2%, respectively, from the archaeal community in all experimental treatments (Fig. 5). The only abundant *Euryarchaeota* members detected were affiliated to an unclassified genus of the Marine Benthic Group D family (MBGD and DHVEG-1), which accounted for 18 to 23% of the archaeal biota in all treatments. Outside of the *Euryarchaeota* phylum, members from the newly named *Bathyarchaeota* lineage (formerly known as Miscellaneous Crenarchaeotic Group) were another cluster of microorganisms that remained in high percentages (from 8 to 14%) in all treatments. Two genera from the *Thaumarchaeota* phylum, one belonging to the
pMC2A209 class, and the other from the Marine Benthic Group B (MBG-B) were also consistently present in all sediment samples showing AOM, the latter one increasing its proportion up to 12% when sulfate reduction was inhibited (Fig. 5). From the bacterial counterpart, the most abundant bacteria in two of the treatments was a genus of *Oceanimonas* from *Aeromonadaceae* family (*Gammaproteobacteria*), whose presence was diminished when sulfate reduction was inhibited and when $^{13}\text{CH}_4$ was absent (Fig. S3), suggesting that this microorganism might have been involved in sulfate-dependent AOM. Other evident changes in the bacterial community included the increase of *Clostridia* and *Bacilli* members when external NOM was supplied (Fig. S3), which agrees with their capacity to reduce HS (28).

**AOM linked to AQDS reduction**

In order to confirm the capacity of the sediment biota to channel $^{13}$C-methane derived electrons to quinone groups, the humic analogue, anthraquinone-2,6-disulfonate (AQDS), was added as an electron acceptor to the artificial basal medium for sediment enrichments. AQDS reduction and methane consumption were observed since the first enrichment cycle, although no clear relationship between net methane consumption and AH$_2$QDS production was observed due to high concentrations of intrinsic electron donors and acceptors (data not shown). Nevertheless, during the third incubation cycle, net AOM was observed within 11 days, which corresponded to a final ratio of oxidized methane/reduced AQDS of 1:4.7 corrected for endogenous controls, which is very close to the stoichiometric 1:4 according to the following equation (Fig. 6a):

$$\text{CH}_4 + \text{AAQDS} + 2\text{H}_2\text{O} \rightarrow \text{CO}_2 + 4\text{AH}_2\text{QDS}$$
Gibbs free energy ($\Delta G^\circ$) = -43.2 kJ mol$^{-1}$

Analysis of 16S rRNA gene sequences from enriched sediment sampled at the end of the third cycle of AQDS-dependent AOM activity (Fig. S1) displayed a significant decrease on the diversity of the microbial community evidenced by a decrease in Shannon index, from 5.52 in freshly sampled sediment to 3.56 after enrichment with CH$_4$ and AQDS. Significant increments and decreases of specific groups of archaea and bacteria did occur in this enrichment (Fig. 6b and c). From the archaeal fraction, the pMC2A209 class from the Thaumarchaeota and the Methanoseta genera were archaeal clusters that significantly increased their presence in the AQDS enrichment (34% and 23%, respectively). Also in the AQDS enrichment, the Bathyarchaeota phylum previously detected in wetland sediments, both in the presence and in the absence of external NOM, significantly increased its proportion in the archaeal community (around 10% respect to the original composition), suggesting potential metabolic arrangements to thrive under AQDS-dependent AOM conditions (Fig. 6b). Humus-reducing bacteria that proliferated throughout the five months of enrichment included genera from the Desulfuromonadales (29, 30), Clostridiales (14, 28) and Propionibacteriales (31) orders in 27%, 7%, and 12%, respectively, with respect to the original composition (Fig. S3).

Discussion

NOM as terminal electron acceptor fueling AOM in wetland sediment. Although the complex composition of the studied wetland sediment challenged efforts to elucidate the microbial processes responsible for the high methanotrophic activities quantified, the...
present study provides multiple lines of evidence demonstrating that electron-accepting functional groups present in its NOM fueled AOM by serving as terminal electron acceptor. Indeed, while sulfate reduction was the predominant process accounting for up to 42.5% of AOM activities, microbial reduction of NOM concomitantly occurred. Furthermore, enrichment of wetland sediment with external NOM, as Pahokee Peat HS, significantly promoted AOM with a quantified amount of \( \sim 100 \text{ nmol } ^{13}\text{C}-\text{CH}_4 \text{ oxidized cm}^{-3} \text{ d}^{-1} \) attributed to this microbial process. Spectroscopic evidence also demonstrated that quinone moieties, main redox functional groups in HS (19), were heterogeneously distributed in the studied wetland sediment and that their reduction occurred during the course of AOM. Moreover, an enrichment derived from wetland sediments performing AOM linked to NOM reduction stoichiometrically oxidized methane coupled to AQDS. Sediment incubations performed in the presence of the sulfate reduction inhibitor, molybdate, further confirmed the role of HS in AOM. Certainly, even though sulfate-reducing activities significantly decreased in the presence of molybdate, AOM activities remained high, while microbial reduction of NOM was doubled under these conditions. These interesting findings suggest that methanotrophic microorganisms performing sulfate-dependent AOM might have directed electrons derived from AOM towards NOM when sulfate reduction became blocked as has been suggested based on experiments performed under artificial conditions (27).

**Microbial communities in wetland sediments performing AOM.** Archaeal clusters consistently found in wetland sediment incubations performing AOM included members from the MBG-D family, which have already been proposed as players in metal-dependent AOM (6), thus their presence agrees with evidence indicating AOM linked to iron.
reduction observed in some experimental controls (Table S2). Additionally, these microorganisms were not found in the AQDS enrichment, probably due to depletion of intrinsic ferric iron throughout the incubation cycles. Archaea constantly present amongst fresh sediment incubation and AQDS enrichment were those from the pMC2A209 class and the Batharchaeota phylum. To our knowledge, the pMC2A209 class of archaea has not been related to AOM, but its close partners from the MBG-B class have been consistently found in environments in which AOM occurs (32–35). In fact, Thaumarchaeota members, including the MBG-B, have been found in consortia performing AOM in the absence of ANME clades (36). Interestingly, the pMC2A209 cluster seemed to duplicate its proportion up to 12% when sulfate reduction was inhibited (by molybdate), which might suggest that the impediment of sulfate reduction enhanced its activity promoting AOM coupled to NOM reduction. Respect to the Batharchaeota phylum, increasing evidence suggests that this lineage might be involved in the methane cycle. Recently, it has been demonstrated that this cluster possesses the necessary genetic elements to express the enzymatic machinery required for methane production, and potentially methane consumption (37). Additionally, Saxton and colleagues have found abundant Batharchaeota representation in a fulvic acids rich deep sediment that oxidizes methane uncoupled from sulfate reduction (22). Unexpectedly, a very low percentage within the archaeal population was identified as members from the ANME type archaea, even though it would be expected to find ANME-2 members since it is the only ANME subgroup with proven capability to derive electrons extracellularly towards humus and its analogues under artificial conditions (27). Our microcosms, both in fresh sediment as well as in the AQDS long-term enrichment, showed a barely detectable number of copies of
ANME-1b and ANME-3 sequences retrieved by the methodology employed, suggesting a low presence of ANME microorganisms in the ecosystem studied.

Regarding the bacterial composition, while *Clostridia*, *Bacilli* and *Gammaproteobacteria* were significantly represented within the fresh sediment performing AOM (Fig. S3), the AQDS enrichment (Fig. 6) exhibited the most significant increase in *Deltaproteobacteria* of the *Desulfuromonadales* order, which includes several humus-reducing microorganisms (14). Since a wide diversity of microorganisms have been proven to reduce humus analogues or HS, we do not rule out that diverse bacterial clusters could have participated in partnership with detected archaea to jointly performed AOM coupled to NOM reduction. Nevertheless, humus-reducing bacteria possess metabolic versatility and capability to reduce miscellaneous electron acceptors, which makes it difficult to come to conclusions about their participation in our experiments. Further investigation must be done to unravel the potential involvement of humus-reducing bacteria in AOM.

**Ecological significance.** To our knowledge this is the first report of AOM coupled to microbial reduction of NOM, which constitutes a missing link within the carbon cycle. HS frequently contribute up to 80% of soil NOM and up to 50% of dissolved NOM in aquatic environments. While the labile fraction of NOM promotes methanogenesis in anaerobic environments, the slowly decomposing humic portion may serve as an important barricade to prevent methane emissions in organotrophic ecosystems by serving as terminal electron acceptor driving AOM (Fig. 7). As an example, considering the maximum AOM driven by microbial reduction of NOM measured in humic enriched sediments, and the global area of coastal wetlands (38, 39), we approximate that this microbial process consumes up to 114 Tg CH₄ yr⁻¹. Considering the global wetland area (10), we anticipate methane suppression
of more than 1,300 Tg yr\(^{-1}\) (see Supplemental Material for details). Accordingly, NOM-driven AOM may be more prominent in organotrophic sites with poor sulfate content, such as peatlands, swamps and organotrophic lakes. This premise is supported by suppression of methanogenesis by HS observed in different ecosystems (16, 17) and by the widespread AOM activity reported across many peatland types (40–42). The potential role of HS is further emphasized because their electron accepting capacity is fully recycled in recurrently anoxic environments. Thus, the suppression of methanogenesis by HS estimated to be of the order of 190,000 mol CH\(_4\) km\(^{-2}\) yr\(^{-1}\) may be much larger than previously considered (43).

Materials and Methods

Sediment sampling and characterization

Sediment cores were collected from the tropical marsh \textit{Sisal}, located in Yucatán Peninsula, south-eastern Mexico (21°09′26″N, 90°03′09″W) in January 2016. Sediment cores with a depth of 15 cm were collected under a water column of approximately 70 cm. Water samples were also collected from the area of sediment sampling points to be used as liquid medium in anaerobic incubations. All sediment and water samples were sealed in hermetic flasks and were maintained in ice until arrival to the laboratory. Upon arrival, all sampled materials were stored at 4°C in a dark room until analysis and incubation. Sediment cores were opened and homogenized within an anaerobic chamber (atmosphere composed of N\(_2\)/H\(_2\) (95%/5%, v/v)) before characterization and incubation. No amendments (addition of chemicals, washing or exposure to air) were allowed on sediment and water samples in
order to reflect the actual conditions prevailing in situ as closely as possible. Characterization of water and sediment samples is described in Table S1.

Sediment incubations. Water samples collected from sediment sampling points were thoroughly mixed before amendment with HS (2.5 g l\(^{-1}\)) by magnetic stirring. *Pahokee Peat* (Florida, Everglades) HS, purchased from the International Humic Substances Society, were employed as external NOM in sediment incubations. Humic-enriched water was flushed with N\(_2\) to blow away any dissolved oxygen. Portions of 15 ml were then distributed in 25-ml serological flasks. Sediment containers were opened inside an anaerobic chamber. Portions of 2.5 ml of wet sediment previously homogenized were then inoculated into each serological bottle. After sealing all bottles with rubber stoppers and aluminum rings inside the anaerobic chamber, they were flushed with N\(_2\). Once anaerobic conditions were established, 5 ml of \(^{13}\)C-labeled methane were injected into each vial to reach a \(^{13}\)CH\(_4\) partial pressure of 0.67 atm in a headspace of 7.5 ml. Controls incubated in the absence of external HS were also prepared by following an identical protocol. Killed controls included chloroform at a concentration of 10\% (v/v) to annihilate any microbial activity. Additional incubations were executed in the presence of the sulfate-reduction inhibitor, molybdate (25 mM), in the presence and in the absence of external NOM. All incubation bottles were statically placed in a dark room at 28 °C (temperature prevailing at *Sisal* wetland at the sampling time). The pH remained at 7.5±0.05 throughout all incubations.

Enrichment incubations with AQDS. Incubations were commenced by inoculating 120-ml serological bottles with 10 g of volatile suspended solids (VSS) l\(^{-1}\) of Sisal sediment.
Prior inoculation, portions of 60 ml of artificial medium were distributed into the incubation bottles and flushed for 15 min with a mixture of N₂:CO₂ (80%/20%, v/v) for stripping any dissolved oxygen from the medium. AQDS (>98.0% purity, TCI AMERICA Chemicals) was added at a concentration of 10 mM as terminal electron acceptor along with the following basal medium components (g l⁻¹): NaHCO₃ (5), NH₄Cl (0.3), K₂HPO₄ (0.2), MgCl₂·6H₂O (0.03) and CaCl₂ (0.1). Trace elements were included in the medium by adding 1 ml l⁻¹ of a solution with the following composition (mg l⁻¹): FeCl₂·4H₂O (2,000), H₂BO₃ (50), ZnCl₂ (50), CuCl₂·6H₂O (90), MnCl₂·4H₂O (500), AlCl₃·6H₂O (90), CoCl₂·6H₂O (2000), NiCl₂·6H₂O (920), Na₂SeO₃·5H₂O (162), (NH₄)₆Mo₇O₂₄ (500), EDTA (1,000), Na₂WO₄·H₂O (100) and 1 ml l⁻¹ of HCl at 36%. The final pH of the medium was 7.2 and no changes were observed throughout the incubation time. Once inoculation took place, microcosms were sealed with rubber stoppers and aluminum rings, and then flushed with the same N₂:CO₂ mixture. After anoxic conditions were established, 1 ml of sodium sulfide stock solution was injected into each vial to reach a sulfide concentration of 0.1 g l⁻¹ in order to consume any traces of dissolved oxygen. Methane was provided into the microcosms by injecting 30 ml of CH₄ (99.9% purity, Praxair) reaching a partial pressure of methane of 0.54 atm. Subsequent incubations were performed after AQDS was reduced (converted to AH₂QDS) coupled to anaerobic oxidation of methane (AOM). A new set of bottles containing basal medium with AQDS (10 mM) were inoculated within an anaerobic chamber by transferring 10 ml of slurry (sediment and medium) taken from previous incubations (Fig. S1). The following incubations were completed under the same experimental conditions.

**Analytical techniques**
Isotopic carbon dioxide and methane measurements. Ions $^{16}$ (12CH$_4$), 17 (13CH$_4$), 44 (12CO$_2$) and 45 (13CO$_2$) were detected and quantified in a Gas Chromatograph Agilent Technologies 7890A coupled to a Mass Spectrometer (detector) Agilent Technologies 5975C, the ionization was achieved by electronic impact and quadrupole analyzer. For the analysis, a capillary column Agilent Technologies HP-PLOT/Q with a stationary phase of poly-styrene-di-vinyl-benzene (30 m × 0.320 mm × 20 µm) was employed as stationary phase using helium as carrier gas. The chromatographic method was as follows: the starting temperature was 70 °C which was held for 3 min, and then a ramp with an increase of 20 °C per min was implemented until 250 °C was reached and maintained for 1 min. The total time of the run had a duration of 13 min. The temperature of the injection port was 250 °C. The injection volume was 20 µl and there was only one replicate of injection per bottle. The gas injected into the GC was taken directly from the headspace of the incubations and immediately injected into the GC port. Methane calibration curves were made by injection of different methane (99.9% of purity) volumes into serological bottles under the same experimental conditions (atmosphere composition, pressure, temperature, and liquid volume). 12CO$_2$ and 13CO$_2$ curves were made using different dried sodium bicarbonate (99% purity, Sigma Aldrich) and sodium 13C-labelled carbonate (99 atom %13C, Sigma Aldrich) concentrations, respectively, in serological bottles which contained the same volume of wetland sediment and water used in incubations. Standards were incubated at room temperature for 12 hours until equilibrium with the gaseous phase was reached. The linear regression analysis of obtained measurements had a co-relation coefficient higher than 0.97. 13CO$_2$ production rates were based on the maximum slope observed on linear regressions considering at least three sampling points.
Methane quantification in AQDS enrichment. Net methane consumption was assessed in terms of methane concentration measurements in the headspace of microcosms. These measurements were carried out by injecting 100 µl of gas samples from the headspace of incubation bottles into a gas chromatograph (Agilent Technologies 6890M) equipped with a thermal conductivity detector, and a Hayesep D (Alltech, Deerfield, Illinois, USA) column with the following dimensions: 3.048 m × 3.185 m × 2.16 mm. Helium was employed as carrier gas at a flux of 12 ml min⁻¹. The temperature of injection port, oven and detector was 250, 60 and 250 °C, respectively. Calibration curves were made for each reaction volume used by injecting different methane concentrations into serological bottles under the same experimental conditions at which microcosms were performed (atmosphere composition, pressure, temperature, and liquid volume).

Determination of electron accepting functional groups in solid phase by XPS. Sediment samples (solid fraction of microcosms) were dried under a constant nitrogen flow after incubation with methane. Once sediments became dried, bottles were open inside an anaerobic chamber with an atmosphere composed of N₂/H₂ (95%/5%, v/v) and were triturated on an agate mortar. Samples were then kept under anaerobic conditions until analysis in a X-Ray Photoelectron Spectroscopy Analyzer PHI VersaProbe II (Physical Electronics, ULVAC-PHI). Two representative spectra were recorded per scanned sample.

Determination of electron accepting functional groups in solid phase by Micro-ATR-FT-IR imaging. Micro-ATR-FT-IR images were collected from each sample with a continuous scan spectrometer, Agilent 660 FT-IR interfaced to a 620 infrared microscope with a 32 × 32 FPA detector and Ge ATR objective for micro-ATR. Each pixel obtains a full IR spectrum or a total of 1024 spectra. Background spectra were collected from a clean
ATR crystal (i.e., without sample). The Ge crystal of the ATR microscope was lowered onto the surface of each sample for a contact area of approximately 100 × 100 μm. Spectra were collected by co-addition of 256 scans over a spectral range of 4000 to 900 cm⁻¹, at a spectral resolution of 4 cm⁻¹. In all images, a color scale bar is set within the software to reflect the relative concentration range, from low to high. Agilent Resolutions Pro was used for data acquisition and analysis.

Determination of electron accepting functional groups in liquid phase by high resolution UV-Vis-NIR spectroscopy. After each incubation cycle, liquid samples (1.5 ml) were taken in an anaerobic chamber with a disposable syringe and put into a quartz cell, which was sealed with plastic film in order to keep anoxic conditions during spectrometric analysis. Spectra were obtained in a Varian Cary 5000 UV–Vis (diffuse reflectance) spectrophotometer, equipped with an integrating sphere.

Nitrite and nitrate determinations. Nitrite and nitrate concentrations were measured according to spectrometric techniques established at Standard Methods (44). Nitrate measurement is taken under acidic conditions at a wavelength of 275 nm and the value obtained is corrected for dissolved organic matter which has its maximum absorbance at 220 nm. Nitrite forms a purple complex through a reaction with sulfanilamide and N-(1-naphthyl) ethylene diamine, which presents its maximum absorbance at a wavelength of 543 nm. Samples were taken with a disposable syringe directly from the microcosms, injected into sealed quartz cuvettes or glass tubes (depending on the required lecture wavelength) and immediately taken to the spectrophotometer to avoid any reaction of the sample with atmospheric oxygen.
Sulfate and sulfide determinations. Samples were extracted from microcosms and immediately filtered through 0.22 µm nitrocellulose membranes. Filtered samples were then diluted (1:10) with deionized water and processed in an Agilent Capillary Electrophoresis System (Agilent Technologies) according to the methodology proposed by Soga & Ross (45). Dissolved sulfide was measured by the spectrometric method proposed by Cord-Ruwisch (46). Briefly, 100 µl of sample were taken and immediately mixed in vortex with 4 ml of an acidic CuSO₄ solution. Absorbance at 480 nm was immediately registered in a UV-VIS spectrophotometer (Thermo Spectronic) to avoid sulfide oxidation before measurements.

Humic substances reduction and ferrous iron measurements. Quantification of the reduction of electron-accepting functional groups in HS was performed according to Lovley et al. (18). Slurry samples (~500 µl) were taken from microcosms with a disposable syringe while bottles were being manually shaken inside an anaerobic chamber. A portion of each sample (200 µl) was mixed with an equal volume of an acidic solution (HCl, 0.5 M) and allowed to stand for 30 min, while the same volume of sample was reacted with ferric citrate (20 mM) for 3 hours. After reaction with ferric citrate, samples were mildly re-suspended in a vortex and 200 µl were left repose with the same volume of HCl solution for 30 min. Afterwards, each sample was centrifuged for 10 min at 10,000 g in a centrifuge. Spectrafuge 16M and 200 µl of supernatant were then recovered and reacted with a solution of 0.2 g l⁻¹ of 2,4,6-tris(2-pyridil)-1,3,5-triazine (ferrozine reagent). Ferrous iron produced due to chemical reduction of ferric citrate by reduced functional groups in HS, forms a purple complex along with ferrozine reagent, which has its maximum absorbance at 562 nm. The ferrozine solution was buffered with HEPES (50 mM). Once centrifuged samples were
mixed with ferrozine solution, they were left reacting for 10 min before their measurement in a spectrometer Thermo Scientific Genesis 10 UV located inside an anaerobic chamber. All solutions employed in this determination were bubbled with N₂ for 30 min to ensure the absence of dissolved oxygen.

**Total carbon (TC), total organic carbon (TOC) and total inorganic carbon (TIC) measurements.** Water samples were filtered through 0.22 µm nitrocellulose membranes and diluted with deionized water, while sediment samples were dried until constant weight. Both liquid and solid samples were analyzed in a Total Organic Carbon analyzer Shimadzu TOCVCS/TNM-1 equipped with a solids sampling port (SSM-5000A). Solid sample processing time was 6 min at 900°C using O₂ (500 ml min⁻¹) with a purity of 99.9% as carrier gas, all samples were analyzed by triplicate.

**Total, volatile and fixed solids.** Total, fixed and volatile solids were measured by triplicate according to Standard Methods procedure (44).

**Elemental composition.** Elemental composition of sediments was assessed by analyzing acid-extracts from 2 g of wet sediment. In the case of iron and manganese measurements in microcosms, supernatant samples were taken with disposable syringes, filtered and acidified prior analysis. Samples were then analyzed by inductively coupled plasma optical emission spectrometry (ICP-OES) in an equipment Varian 730-ES. The operational conditions were: potency 1 kW, auxiliary flow: 1.5 l min⁻¹, net flow: 0.75 l min⁻¹, sample taking delay: 30 s, and the number of measured replicates by sample was three. Argon was employed as carrier gas.
**DNA extraction, PCR amplification and sequencing.** One microcosm for each selected treatment was randomly chosen at the end of the incubation period (30 days for experiments presented in Fig. 1, and 151 days for experiments depicted in Fig. 6). Before DNA extraction, liquid medium was decanted and extracted from the serological bottles. The total sediment was homogenized afterwards and a subsample of 0.5 g was taken to proceed with DNA extraction. The remaining sediment and the other microcosms were used for material characterization. The total DNA was extracted from sediment samples using the Power Soil DNA extraction kit (MO BIO Laboratories, Carlsbad, CA, USA) following the protocol described by the manufacturer. DNA isolated from each sample was amplified using primers 341F and 785R, targeting the V3 and V4 regions of the 16S rRNA gene fused with Illumina adapter overhang nucleotide sequences (47). The polymerase chain reactions (PCRs) were performed in 50 μl reactions using Phusion Taq polymerase (Thermo Scientific, USA) under the following conditions: denaturation at 98 °C for 60 s, followed by 5 cycles of amplification at 98 °C for 60 s, 50 °C for 30 s and 72 °C for 30 s, followed by 25 cycles of amplification at 98 °C for 60 s, 55 °C for 30 s and 72 °C for 30 s, followed by a final extension of 72 °C for 5 min. Two independent PCR reactions were performed for each sample. The products were indexed using Illumina’s 16S Metagenomic Sequencing Library Preparation protocol and Nextera XT Index Kit v2 (Illumina, San Diego CA). Libraries were deep sequenced with the Illumina MiSeq sequencer.

**Bioinformatics Analysis.** An analysis of 16S rRNA gene libraries was carried out using Mothur open source software package (v 1.34.4) (48). The high quality sequence data were analyzed for potential chimeric reads using the UCHIME algorithm. Sequences containing homopolymer runs of 9 or more bases, those with more than one mismatch to the
sequencing primer and Q-value average below 25 were eliminated. Group membership was
determined prior to the trimming of the barcode and primer sequence. Sequences were
aligned against the SILVA 123 16S/18S rRNA gene template using the nearest alignment
space termination (NAST) algorithm, and trimmed for the optimal alignment region. A
pairwise distance matrix was calculated across the non-redundant sequence set, and reads
were clustered into operational taxonomic units (OTUs) at 3% distance using the furthest
neighbor method. The sequences and OTUs were categorized taxonomically using
Mothur’s Bayesian classifier and the SILVA 123 reference set. The sequences obtained
have been submitted to NCBI GeneBank database.

Accession numbers

The accession numbers of sequences in this work were deposited in the GenBank sequence
read archive under the BioProject with SRP094593 accession number.

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FIGURE LEGENDS

Fig. 1. Anaerobic methane oxidation measured as $^{13}\text{C} \text{CO}_2$ production in microcosms’ headspace and $^{13}\text{C}$ enrichment calculated as $^{13}\text{F}_{\text{CO}_2}$ ($^{13}\text{CO}_2/\left(^{13}\text{CO}_2 + ^{12}\text{CO}_2\right)$). Panel a displays the kinetics for incubations performed with unamended sediment. Panel b displays the kinetics for incubations performed with sediment enriched with 2.5 g l$^{-1}$ of external NOM in the form of Pahokee Peat humic substances. Error bars represent the standard error among replicates ($n = 4$, or $3^*$). SR-INH stands for sediment incubations performed with molybdate (25 mM) in order to inhibit sulfate reduction. $^{13}\text{CO}_2$ production rates were based on the maximum slope observed on linear regressions considering at least three sampling points.

Fig. 2. Production of $^{13}\text{CO}_2$ and reduction of intrinsic or added electron acceptors at the end of the exponential phase (20 days of incubation) in the absence (Panels a and b) and in the presence (Panels c and d) of external NOM as HS from Pahokee Peat. SR-INH stands for controls amended with sulfate-reduction inhibitor, sodium molybdate (25 mM). Error bars represent the standard error among replicates. $^{13}\text{CO}_2$ produced was measured as described for Fig.1. Quantification of sulfate and nitrate reduction imply decrease on their concentration at this sampling time, whereas Fe(III) reduction was quantified in terms of the ferrous iron produced. Reduction of NOM and HS was determined by the ferrozine technique.
Fig. 3. Spectroscopic evidence of the presence of quinone moieties and their reduction in wetland sediment samples. Panels a and b depict the Micro-ATR-FTIR representative spectra taken from imaged areas generated after processing quinone functional groups (1650-1620 cm\(^{-1}\)) of sediment samples before incubation in the absence and in the presence, respectively, of external NOM in the form of Pahokee Peat HS. Panels c and e portray XPS high resolution profiles of C1s, while d and f represent O1s signal. Panels c and d belong to sediment samples prior incubation, while panels e and f correspond to sediment samples after incubation with \(^{13}\)C-methane. Regions and components were corrected at 284.8 eV for the C-C adventitious carbon A; B and G components belong to C-O bond (\(~286.6\) and \(~532\) eV), C and H correspond to C=O functional group (\(~288.9\) and \(~533.3\) eV), D belongs to –COOH (\(~289.6\) ), E is typical of the presence of carbonate (\(~291\) ) and F suggests the occurrence of a metallic oxide (\(~530\) ).

Fig. 4. High performance Ultraviolet-Visible-Near Infrared spectra obtained from liquid samples before and after incubation with \(^{13}\)CH\(_4\). Panels a and c depict spectra obtained before incubation with \(^{13}\)CH\(_4\), while panels b and d show spectra obtained after incubation with \(^{13}\)CH\(_4\).

Fig. 5. Archaeal composition in wetland sediment samples performing AOM. Most abundant archaeal genera detected, based on 16S rRNA amplicon gene libraries, on
selected experimental treatments shown in Fig. 1 at the end of the incubation period (30 days).

Fig. 6. AOM with AQDS as electron acceptor by an enrichment derived from wetland sediment. Panel a: Kinetics of methane consumption linked to AQDS reduction (to \( \text{AH}_2\text{QDS} \)) observed during the last 11 days of the entire enrichment process lasting 151 days: filled squares (-■-) represent microcosms including \( \text{CH}_4 \) as electron donor and AQDS as electron acceptor (complete experiments, \( n=3 \)), open squares (-□-) represent controls without electron acceptor provided (without AQDS control, \( n=3 \)), solid circles (-●-) represent \( \text{CH}_4 \) free microcosms (endogenous controls, \( n=3 \)), and crosses (-×-) represent heat killed controls (sterile controls, \( n=2 \)). Error bars represent the standard error among replicates. Panels b and c depict microbial community changes at the end of the enrichment (151 days of incubation) at the phylum level based on Illumina sequencing of 16S rRNA V3-V4 regions. Fresh sediment composition was used as a reference.

Figure 7. Schematic representation of methane generation and consumption by wetland sediment biota. While a fraction of NOM may serve as electron acceptor to support AOM (NOM-AOM) and decouple sulfate-reduction dependent AOM (SR-AOM), depending on its chemical properties, a labile fraction of NOM could also be degraded following the methanogenesis pathway by a fermenting and methanogenic fraction of the consortia. Equilibrium between these three phenomena must be tightly dependent on thermodynamic conditions, concentration of chemical species, and composition of
microbial community. *Anaerobic methanotrophic archaea are considered in a broader perspective than ANME clades from *Euryarchaeota* phylum.
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**Abundance code**